# CONSORT-EHEALTH (V 1.6.1) - Submission/Publication Form

The CONSORT-EHEALTH checklist is intended for authors of randomized trials evaluating web-based and Internet-based applications/interventions, including mobile interventions, electronic games (incl multiplayer games), social media, certain telehealth applications, and other interactive and/or networked electronic applications. Some of the items (e.g. all subitems under item 5 - description of the intervention) may also be applicable for other study designs.

The goal of the CONSORT EHEALTH checklist and guideline is to be

- a) a guide for reporting for authors of RCTs,
- b) to form a basis for appraisal of an ehealth trial (in terms of validity)

CONSORT-EHEALTH items/subitems are MANDATORY reporting items for studies published in the Journal of Medical Internet Research and other journals / scientific societies endorsing the checklist.

Items numbered 1., 2., 3., 4a., 4b etc are original CONSORT or CONSORT-NPT (non-pharmacologic treatment) items. Items with Roman numerals (i., ii, iii, iv etc.) are CONSORT-EHEALTH extensions/clarifications.

As the CONSORT-EHEALTH checklist is still considered in a formative stage, we would ask that you also RATE ON A SCALE OF 1-5 how important/useful you feel each item is FOR THE PURPOSE OF THE CHECKLIST and reporting quideline (optional).

Mandatory reporting items are marked with a red \*.

In the textboxes, either copy & paste the relevant sections from your manuscript into this form - please include any quotes from your manuscript in QUOTATION MARKS,

or answer directly by providing additional information not in the manuscript, or elaborating on why the item was not relevant for this study.

YOUR ANSWERS WILL BE PUBLISHED AS A SUPPLEMENTARY FILE TO YOUR PUBLICATION IN JMIR AND ARE CONSIDERED PART OF YOUR PUBLICATION (IF ACCEPTED).

Please fill in these questions diligently. Information will not be copyedited, so please use proper spelling and grammar, use correct capitalization, and avoid abbreviations.

DO NOT FORGET TO SAVE AS PDF \_AND\_ CLICK THE SUBMIT BUTTON SO YOUR ANSWERS ARE IN OUR DATABASE !!!

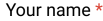
Citation Suggestion (if you append the pdf as Appendix we suggest to cite this paper in the caption): Eysenbach G, CONSORT-EHEALTH Group

CONSORT-EHEALTH: Improving and Standardizing Evaluation Reports of Web-based and Mobile Health Interventions J Med Internet Res 2011;13(4):e126

URL: <a href="http://www.jmir.org/2011/4/e126/">http://www.jmir.org/2011/4/e126/</a>

doi: 10.2196/jmir.1923 PMID: 22209829

\* Required



Primary Affiliation (short), City, Country University of Toronto, Toronto, Canada	*
Duke University, Durham, NC	
Your e-mail address *  abc@gmail.com	
michele.patel@stanford.edu	
Title of your manuscript * Provide the (draft) title of your manuscript.	
Comparing self-monitoring strategies for weight controlled trial	loss in a smartphone app: A randomized
Name of your App/Software/Interventi If there is a short and a long/alternate name, write the short GoalTracker	
If there is a short and a long/alternate name, write the shor	

# URL of your Intervention Website or App

e.g. a direct link to the mobile app on app in appstore (itunes, Google Play), or URL of the website. If the intervention is a DVD or hardware, you can also link to an Amazon page.

URL of an image/screenshot (optional)
Your answer
Accessibility * Can an enduser access the intervention presently?
access is free and open
access only for special usergroups, not open
access is open to everyone, but requires payment/subscription/in-app purchases
app/intervention no longer accessible
Other: utilized a commercial mobile app for self-monitoring (MyFitnessPal), which is freely
Primary Medical Indication/Disease/Condition * e.g. "Stress", "Diabetes", or define the target group in brackets after the condition, e.g. "Autism (Parents of children with)", "Alzheimers (Informal Caregivers of)"
Obesity
Primary Outcomes measured in trial * comma-separated list of primary outcomes reported in the trial  Weight (kg)
Secondary/other outcomes  Are there any other outcomes the intervention is expected to affect?
Intervention engagement

# Recommended "Dose" \*

What do the instructions for users say on how often the app should be used?

Approximately Weekly Approximately Monthly Approximately Yearly "as needed" Other:  Approx. Percentage of Users (starters) still using the app as recommended after 3 months * unknown / not evaluated 0-10% 11-20% 21-30% 31-40% 41-50% 51-60% 61-70% 6 71%-80% 91-100% Other:	Approximately Daily
<ul> <li>Approximately Yearly</li> <li>"as needed"</li> <li>Other:</li> <li>Approx. Percentage of Users (starters) still using the app as recommended after 3 months *</li> <li>unknown / not evaluated</li> <li>0-10%</li> <li>11-20%</li> <li>21-30%</li> <li>31-40%</li> <li>41-50%</li> <li>51-60%</li> <li>61-70%</li> <li>71%-80%</li> <li>81-90%</li> <li>91-100%</li> </ul>	Approximately Weekly
<ul> <li>○ "as needed"</li> <li>○ Other:</li> <li>Approx. Percentage of Users (starters) still using the app as recommended after 3 months *</li> <li>○ unknown / not evaluated</li> <li>○ 0-10%</li> <li>○ 11-20%</li> <li>○ 21-30%</li> <li>○ 31-40%</li> <li>○ 41-50%</li> <li>○ 51-60%</li> <li>○ 61-70%</li> <li>○ 71%-80%</li> <li>○ 81-90%</li> <li>○ 91-100%</li> </ul>	Approximately Monthly
Other:  Approx. Percentage of Users (starters) still using the app as recommended after 3 months *  unknown / not evaluated  0-10%  11-20%  21-30%  31-40%  41-50%  51-60%  61-70%  71%-80%  81-90%  91-100%	Approximately Yearly
Approx. Percentage of Users (starters) still using the app as recommended after 3 months *  unknown / not evaluated  0-10%  11-20%  21-30%  31-40%  41-50%  51-60%  61-70%  71%-80%  81-90%  91-100%	○ "as needed"
after 3 months *  unknown / not evaluated  0-10%  11-20%  21-30%  31-40%  41-50%  51-60%  61-70%  71%-80%  91-100%	Other:
<ul> <li>○ 0-10%</li> <li>○ 11-20%</li> <li>○ 21-30%</li> <li>○ 31-40%</li> <li>○ 41-50%</li> <li>○ 51-60%</li> <li>○ 61-70%</li> <li>○ 71%-80%</li> <li>○ 81-90%</li> <li>○ 91-100%</li> </ul>	
<ul> <li>☐ 11-20%</li> <li>☐ 21-30%</li> <li>☐ 31-40%</li> <li>☐ 41-50%</li> <li>☐ 51-60%</li> <li>☐ 61-70%</li> <li>⑥ 71%-80%</li> <li>☐ 81-90%</li> <li>☐ 91-100%</li> </ul>	unknown / not evaluated
<ul> <li>21-30%</li> <li>31-40%</li> <li>41-50%</li> <li>51-60%</li> <li>61-70%</li> <li>71%-80%</li> <li>81-90%</li> <li>91-100%</li> </ul>	O-10%
<ul> <li>○ 31-40%</li> <li>○ 41-50%</li> <li>○ 51-60%</li> <li>○ 61-70%</li> <li>○ 71%-80%</li> <li>○ 81-90%</li> <li>○ 91-100%</li> </ul>	O 11-20%
<ul> <li>↓ 41-50%</li> <li>↓ 51-60%</li> <li>♠ 61-70%</li> <li>♠ 71%-80%</li> <li>♠ 81-90%</li> <li>♠ 91-100%</li> </ul>	O 21-30%
<ul> <li>51-60%</li> <li>61-70%</li> <li>71%-80%</li> <li>81-90%</li> <li>91-100%</li> </ul>	31-40%
<ul> <li>61-70%</li> <li>71%-80%</li> <li>81-90%</li> <li>91-100%</li> </ul>	O 41-50%
<ul><li>71%-80%</li><li>81-90%</li><li>91-100%</li></ul>	<u></u>
<ul><li>81-90%</li><li>91-100%</li></ul>	O 61-70%
91-100%	71%-80%
	81-90%
Other:	91-100%
	Other:

Overall, was the app/intervention effective? \*

yes: all primary outcomes were significantly better in intervention group vs control

partly: SOME primary outcomes were significantly better in intervention group vs control
no statistically significant difference between control and intervention
outcomes potentially harmful: control was significantly better than intervention in one or more
inconclusive: more research is needed
Other:
Article Preparation Status/Stage *  At which stage in your article preparation are you currently (at the time you fill in this form)
onot submitted yet - in early draft status
ont submitted yet - in late draft status, just before submission
submitted to a journal but not reviewed yet
submitted to a journal and after receiving initial reviewer comments
submitted to a journal and accepted, but not published yet
O published
Other:

onot submitted yet / unclear where I will submit this
<ul><li>Journal of Medical Internet Research (JMIR)</li></ul>
JMIR mHealth and UHealth
JMIR Serious Games
JMIR Mental Health
JMIR Public Health
JMIR Formative Research
Other JMIR sister journal
Other:
Is this a full powered effectiveness trial or a pilot/feasibility trial? *  Pilot/feasibility  Fully powered
Manuscript tracking number *  If this is a JMIR submission, please provide the manuscript tracking number under "other" (The ms tracking number can be found in the submission acknowledgement email, or when you login as author in JMIR. If the paper is already published in JMIR, then the ms tracking number is the four-digit number at the end of the DOI, to be found at the bottom of each published article in JMIR)  on ms number (yet) / not (yet) submitted to / published in JMIR  Other:
TITLE AND ABSTRACT
1a) TITLE: Identification as a randomized trial in the title

# 1a) Does your paper address CONSORT item 1a? \* I.e does the title contain the phrase "Randomized Controlled Trial"? (if not, explain the reason under "other")

( yes								
1a-i) Identify the mode of delivery in the title  Identify the mode of delivery. Preferably use "web-based" and/or "mobile" and/or "electronic game" in the title. Avoid ambiguous terms like "online", "virtual", "interactive". Use "Internet-based" only if Intervention includes non-web-based Internet components (e.g. email), use "computer-based" or "electronic" only if offline products are used. Use "virtual" only in the context of "virtual reality" (3-D worlds). Use "online" only in the context of "online support groups".  Complement or substitute product names with broader terms for the class of products (such as "mobile" or "smart phone" instead of "iphone"), especially if the application runs on different platforms.								
	1	2	3	4	5			
subitem not at all important	0	0	0	0	0	essential		
Copy and paste relevant s quotes from your manusc explain why the item is no "Comparing self-mor	cript), or elab ot applicable,	orate on this ite relevant for yo	em by providir ur study	ng additional in	formation not i			
1a-ii) Non-web-b Mention non-web-based o		•	•					
	1	2	3	4	5			
subitem not at all important	0	0	0	0	0	essential		
Does your paper Copy and paste relevant s quotes from your manusc explain why the item is no	sections fron cript), or elab ot applicable,	n manuscript tit orate on this ite /relevant for yo	tle (include qu em by providir ur study	ng additional in	formation not i			

# 1a-iii) Primary condition or target group in the title

	1	2	3	4	5					
subitem not at all important	0	0	0	0	0	essential				
Does your paper address subitem 1a-iii? *  Copy and paste relevant sections from manuscript title (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study  "for weight loss"										
1b) ABSTRACT: conclusions	Structure	ed summa	ary of tria	design,	methods, ı	results, and				
NPT extension: Descrip	ition of expe	rimental treat	ment, compa	rator, care pr	oviders, center	s, and blinding				
1b-i) Key feature comparator in the Mention key features/fur mention theories and printindexers by including implication is missing from	ne METH actionalities/onciples used to portant synon	ODS sections of the section of the s	ion of the the interventione site. Keep in	ABSTRA on and compa on mind the need a abstract what	CT rator in the absteds of systemat	tract. If possible, also ic reviewers and				
	1	2	3	4	5					
subitem not at all important	0	0	0	0	0	essential				
Does your paper Copy and paste relevant indicate direct quotes fro ms, or briefly explain why "12-week standalone either (1) both weigh (2) weight through w or (3) only diet (App-	sections from om your manu of the item is not e weight los of and diet, or eek 4, ther	n the manuscrip uscript), or elab ot applicable/r ss interventi with weekly	pt abstract (indorate on this it elevant for you on using the lessons, ac	em by providi ir study e MyFitness tion plans,	ng additional in sPal app for s and feedbac	formation not in the self- monitoring k (Simultaneous);				

# 1b-ii) Level of human involvement in the METHODS section of the ABSTRACT

abotract what the main paper is reporting. If this information is missing from the main body of text, consider adding ity									
	1	2	3	4	5				
subitem not at all important	0	0	0	0	0	essential			
Does your paper address subitem 1b-ii?  Copy and paste relevant sections from the manuscript abstract (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study  "GoalTracker was an automated RCT (n=105) among adults with overweight or obesity "									
1b-iii) Open vs. closed, web-based (self-assessment) vs. face-to-face assessments in the METHODS section of the ABSTRACT  Mention how participants were recruited (online vs. offline), e.g., from an open access website or from a clinic or a closed online user group (closed usergroup trial), and clarify if this was a purely web-based trial, or there were face-to-face components (as part of the intervention or for assessment). Clearly say if outcomes were self-assessed through questionnaires (as common in web-based trials). Note: In traditional offline trials, an open trial (open-label trial) is a type of clinical trial in which both the researchers and participants know which treatment is being administered. To avoid confusion, use "blinded" or "unblinded" to indicated the level of blinding instead of "open", as "open" in web-based trials usually refers to "open access" (i.e. participants can self-enrol). (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it)									
	1	2	3	4	5				
subitem not at all important	0	$\circ$	0	$\bigcirc$	0	essential			
Does your paper address subitem 1b-iii?  Copy and paste relevant sections from the manuscript abstract (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study  "Participants were recruited via on- and off-line methods. Weight was collected in-person at baseline, 1-, and 3 months using calibrated scales and via self-report at 6 months. Other assessment data were collected in-person via self-report questionnaires."									

## 1b-iv) RESULTS section in abstract must contain use data

Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it)									
	1	2	3	4	5				
subitem not at all important	0	0	0	0	0	essential			
Does your paper address subitem 1b-iv?  Copy and paste relevant sections from the manuscript abstract (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study									
"The median [IQR] no (once began), 5.3 [4.				•	t was 1.9 [5.	3] in Sequential			
1b-v) CONCLUSI Conclusions/Discussions outcome not changed), a uptake and discuss reaso missing from the main be	s in abstract fo nd the interver ons. (Note: On	or negative tria ntion was not ly report in the	als: Discuss th used, discuss e abstract wha	e primary outc whether negat	ome - if the tria tive results are	l is negative (primary attributable to lack of			
	1	2	3	4	5				
subitem not at all important	0	0	0	0	0	essential			
Does your paper address subitem 1b-v?  Copy and paste relevant sections from the manuscript abstract (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study  Your answer									
INTRODUCTION									
2a) In INTRODU	CTION: So	cientific b	ackgrour	nd and ex	planation	of rationale			

# 2a-i) Problem and the type of system/solution

Describe the problem and the type of system/solution that is object of the study: intended as stand-alone intervention vs. incorporated in broader health care program? Intended for a particular patient population? Goals of the

nitervention, e.g., being i Details about the interve				еріасе оі соп	ιριειτιειτι στιτει	รบเนเเบกร <i>ะ</i> (เทบเษ.			
	1	2	3	4	5				
subitem not at all important	0	0	0	0	0	essential			
Does your paper address subitem 2a-i? *  Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study  Improvements are needed to increase self-monitoring engagement in weight loss trials:  "Despite its utility, dietary self-monitoring typically declines over the course of treatment.[2-3]									
Novel strategies are involves enriching s	needed to elf-monitor	improve die	etary self-mo er theoretic	onitoring en ally and em	gagement.[4 pirically sup	4] One strategy ported behavior			
change techniques, such as tailored goals and feedback, action plans, and skills training.[5-6] "  We opted to use a standalone digital approach to maximize self-monitoring engagement: "We tested this sequential strategy using a remotely-delivered intervention that utilizes a popular commercially available mobile app — MyFitnessPal. Utilizing technology for self-monitoring dietary intake has been shown to produce greater adherence and less-pronounced declines in engagement than traditional paper- based tracking methods.[14-15] Interventions without counseling that utilize commercial technology for dietary self-monitoring have produced clinically meaningful weight losses between 2.5-5.5 kg at 3 months and beyond in recent studies.[16-20]"									
2a-ii) Scientific background, rationale: What is known about the (type of) system  Scientific background, rationale: What is known about the (type of) system that is the object of the study (be sure to discuss the use of similar systems for other conditions/diagnoses, if appropriate), motivation for the study, i.e. what are the reasons for and what is the context for this specific study, from which stakeholder viewpoint is the study performed, potential impact of findings [2]. Briefly justify the choice of the comparator.									
	1	2	3	4	5				
subitem not at all important	0	0	0	0	0	essential			

# Does your paper address subitem 2a-ii? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study.

"Utilizing technology for self-monitoring dietary intake has been shown to produce greater adherence and less-pronounced declines in engagement than traditional paper-based tracking methods.[14-15] Interventions without counseling that utilize commercial technology for dietary self-monitoring have produced clinically meaningful weight losses between 2.5-5.5 kg at 3 months and beyond in recent studies.[16-20]"

#### 2b) In INTRODUCTION: Specific objectives or hypotheses

#### Does your paper address CONSORT subitem 2b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"In a randomized trial, we tested a weight loss intervention among adults with overweight or obesity that targeted the aforementioned strategies of including empirically supported behavior change techniques, promoting mastery and self- efficacy through self-monitoring of body weight prior to diet, and utilizing a commercially available app (MyFitnessPal) with high acceptability. [21] We hypothesized that a sequential approach would produce greater weight loss and self-monitoring engagement at 3 months, compared to a traditional simultaneous approach and to an "off-the-shelf" app."

#### **METHODS**

3a) Description of trial design (such as parallel, factorial) including allocation ratio

"GoalTracker was a 3-arm randomized controlled trial comparing three standalone weight loss interventions: (1) a Simultaneous self-monitoring arm in which participants simultaneously tracked body weight and dietary intake each day, and received additional empirically-supported behavior change techniques (weekly lessons, action plans, tips, tailored feedback) via email for the entirety of the intervention; (2) a Sequential arm, consisting of identical intervention components, but allowing for mastery of one skill (i.e., self-monitoring of body weight) before beginning self-monitoring of diet; and (3) an App-Only arm that tracked only diet with no additional behavior change components. Study evaluation visits were held at baseline, 1 month, and 3 months"

"Using simple random sampling, participants were then randomized by study staff to one of three treatment arms using Excel's random number generator to allocate participants equally (1:1:1) across conditions"

# 3b) Important changes to methods after trial commencement (such as eligibility criteria), with reasons

#### Does your paper address CONSORT subitem 3b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"To promote generalizability of findings, during the trial, the BMI criteria were expanded to include participants in the 40.0-45.0 kg/m2 range and the weight change criteria were adjusted to no longer exclude individuals who gained more than 10 lbs in the past 6 months."

#### 3b-i) Bug fixes, Downtimes, Content Changes

Bug fixes, Downtimes, Content Changes: ehealth systems are often dynamic systems. A description of changes to methods therefore also includes important changes made on the intervention or comparator during the trial (e.g., major bug fixes or changes in the functionality or content) (5-iii) and other "unexpected events" that may have influenced study design such as staff changes, system failures/downtimes, etc. [2].

	1	2	3	4	5	
subitem not at all important	0	0	0	0	$\circ$	essential

#### Does your paper address subitem 3b-i?

#### 4a) Eligibility criteria for participants

#### Does your paper address CONSORT subitem 4a? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Inclusion criteria comprised men and women ages 21-65 years old with a body mass index (BMI) between 25.0-45.0 kg/m2 who were interested in losing weight through dietary change. We required participants to have an iPhone or Android smartphone, email address, access to a bathroom scale, and written English fluency. We excluded participants if they were enrolled in another weight loss intervention, used MyFitnessPal to track diet in the past 6 months, lost ≥10 lbs or used a weight loss medication in the past 6 months, had previous or planned bariatric surgery, or if weight loss would be contraindicated (e.g., pregnancy, or in need of medical or psychiatric intervention). To promote generalizability of findings, during the trial, the BMI criteria were expanded to include participants in the 40.0-45.0 kg/m2 range and the weight change criteria were adjusted to no longer exclude individuals who gained more than 10 lbs in the past 6 months."

#### 4a-i) Computer / Internet literacy

Computer / Internet literacy is often an implicit "de facto" eligibility criterion - this should be explicitly clarified.

	1	2	3	4	5	
subitem not at all important	$\circ$	0	0	0	0	essential

#### Does your paper address subitem 4a-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

#### 4a-ii) Open vs. closed, web-based vs. face-to-face assessments:

participant. In online-only was possible or whether the detect/prevent these.	trials, clarify	if participants	were quasi-a	nonymous and	whether havin	g multiple identities		
	1	2	3	4	5			
subitem not at all important	0	0	0	0	0	essential		
Does your paper address subitem 4a-ii? *  Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study								
In general, all assess self-report questionn corresponding evalua Qualtrics for action p measured in person	aires or di ation data blans. To e	rect weight/ (outcomes nsure accur	/height mea measureme ate weight	asurement). ent) were co as often as <sub>l</sub>	Intervention llected via s possible, we	content and smartphone app or eight was		
4a-iii) Informatio Information given during consent procedures (e.g., information may have an	recruitment. publish the	Specify how painformed cons	articipants we ent document	re briefed for re ation as appen	dix, see also it	em X26), as this		
subitem not at all important	0	0	0	0	0	essential		
Does your paper address subitem 4a-iii?  Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study								

Your answer

4b) Settings and locations where the data were collected

## Does your paper address CONSORT subitem 4b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study.

"between April-September 2017 in central North Carolina"

#### 4b-i) Report if outcomes were (self-)assessed through online questionnaires

Clearly report if outcomes were (self-)assessed through online questionnaires (as common in web-based trials) or otherwise.

	1	2	3	4	5	
subitem not at all important	$\circ$	0	0	0	0	essential

#### Does your paper address subitem 4b-i? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

To ensure accurate weight as often as possible, weight was measured in person at baseline, 1- and 3-months, and was self-reported at 6 month follow-up. Height was measured in-person at baseline. Engagement data were collected via smartphone app (or Qualtrics for the action plans). All other measures were self-assessed through online questionnaires.

#### 4b-ii) Report how institutional affiliations are displayed

Report how institutional affiliations are displayed to potential participants [on ehealth media], as affiliations with prestigious hospitals or universities may affect volunteer rates, use, and reactions with regards to an intervention.(Not a required item – describe only if this may bias results)

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

#### Does your paper address subitem 4b-ii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

5) The interventions for each group with sufficient details to allow replication, including how and when they were actually administered

# 5-i) Mention names, credential, affiliations of the developers, sponsors, and owners

manuscript).	, tills fleeds	to be decialed	iii a Commet C	or interest sec	tion of mention	ed elsewhere in the
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper Copy and paste relevant s quotes from your manusc explain why the item is no	sections fron cript), or elab	n the manuscri orate on this it	pt (include quo em by providir			
5-ii) Describe the Describe the history/deve usability testing), as these	lopment pro	cess of the ap	olication and p	revious forma		
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper Copy and paste relevant s quotes from your manusc explain why the item is no	sections fron cript), or elab	n the manuscri orate on this it	pt (include que em by providir			
Your answer						
5-iii) Revisions at Revisions and updating. Of comparator, if applicable) evaluation process, or who components such as new (for unexpected events see	Clearly mention evaluated, controlled evaluated, controlled evaluated, controlled evaluated evaluated, controlled evaluated, controlled evaluated evaluated, controlled evaluated evaluate	on the date and or describe whe velopment and	ether the interv or content wa	ention underwas "frozen" duri	ent major chaning the trial. Des	ges during the scribe dynamic
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

# Does your paper address subitem 5-iii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study.

#### Does your paper address subitem 5-v?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

#### 5-vi) Digital preservation

consider creating demo		-	, ,	•	yın süretis G	ninot be archived,
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your pape Copy and paste relevant quotes from your manu explain why the item is	t sections from script), or elabo	the manuscri orate on this it	pt (include qu em by providi	•		
Your answer						
5-vii) Access Access: Describe how por not, whether they had the platform and Internet account or demo mode	l to be a memb et" [1]. To ensur	er of specific re access for e	group. If know editors/review	n, describe hovers/readers, co	w participants ensider to provi	obtained "access to de a "backdoor" login
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your pape Copy and paste relevant quotes from your manu- explain why the item is "All treatment arms that allows users to database with over text format provide	t sections from script), or elabo not applicable/ self-monito o log food ar 5 million foo	the manuscri brate on this it relevant for your red dietary nd beverage ods (myfitne	pt (include qu em by providi our study intake using s, and prov esspal.com	ng additional in g MyFitness ides nutrition ). In-app fee	formation not Pal, a free conal informat	in the ms, or briefly ommercial app ion from a
"During the baseling the MyFitnessPal m		d study sta	ff assiste	ed participar	nts in installi	ng and navigating
MyFitnessPal is fre access were exclud	=	<del>-</del>	with a sma	tphone; indi	viduals with	out smartphone

5-viii) Mode of delivery, features/functionalities/components of the intervention and comparator, and the theoretical framework

theoretical framework [6] used to design them (instructional strategy [1], behaviour change techniques, persuasive features, etc., see e.g., [7, 8] for terminology). This includes an in-depth description of the content (including where it is coming from and who developed it) [1]," whether [and how] it is tailored to individual circumstances and allows users to track their progress and receive feedback" [6]. This also includes a description of communication delivery channels and – if computer-mediated communication is a component – whether communication was synchronous or asynchronous [6]. It also includes information on presentation strategies [1], including page design principles, average amount of text on pages, presence of hyperlinks to other resources, etc. [1].

	1	2	3	4	5	
subitem not at all important	0	0	0	$\circ$	0	essential

#### Does your paper address subitem 5-viii? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

MyFitnessPal was used as designed. Control Theory and Social Cognitive Theory were used to design supplementary intervention content and the schedule of intervention components to include mastery.

"A second strategy includes building mastery, self-efficacy, and self-regulation – key constructs of behavior change in Carver's Control Theory [7] and Bandura's Social Cognitive Theory[8] – before asking participants to engage in dietary self- monitoring. Fostering self-regulatory skills may provide an opportunity for mastery and, in turn, strengthen self-efficacy,[9] which has been linked to greater weight reduction.[10] We propose a novel solution that aims to attenuate the decline in engagement by employing a sequential [11] self-monitoring approach, wherein individuals track only body weight for a period of time and then begin to track diet. Tracking body weight was chosen as it requires minimal effort, provides an opportunity for habit formation (e.g., track every morning upon waking), and is efficacious for weight loss.[12] We focused on only self-monitoring of body weight during the first month, based on prior research demonstrating that enhanced engagement in the first month of treatment may have long-lasting repercussions. [13]"

## 5-ix) Describe use parameters

Describe use parameters (e.g., intended "doses" and optimal timing for use). Clarify what instructions or recommendations were given to the user, e.g., regarding timing, frequency, heaviness of use, if any, or was the intervention used ad libitum.

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

#### Does your paper address subitem 5-ix?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the providing additional information of the ms, or briefly available to the providing additional information of the ms, or briefly available to the providing additional information of the ms, or briefly available to the manuscript of the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the manuscript of the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the manuscript of the

'Self-monitoring of b	ody weigh	t: daily via th	ne app"			
'Self-monitoring of d	ietary intal	ke: daily via	the app"			
5-x) Clarify the lectority the lectority the level of human intervention or as co-interpassistance offered, the tiressistance is delivered. It is and the level of humatem 21 – generalizability.	involvement vention (deta ning and frea t may be nec an involveme	c (care provider ail number and quency of the s essary to disti	es or health pro expertise of p support, how it nguish betwee	rofessionals in is initiated, an In the level of h	nvolved, if any, d the medium numan involve	as well as "type of by which the ment required for the
	1	2	3	4	5	
subitem not at all	$\bigcirc$	$\circ$	0	0	0	essential
Copy and paste relevant s quotes from your manusc explain why the item is no	sections from cript), or elab	n the manuscri orate on this it	pt (include quo em by providin			
Does your paper Copy and paste relevant s quotes from your manusc explain why the item is no	sections from cript), or elab ot applicable,	n the manuscri orate on this it relevant for yo	pt (include quo em by providin our study			
Does your paper Copy and paste relevant sequotes from your manusce explain why the item is not Your answer  5-xi) Report any paper Report any prompts/reminal polication, what triggered prompts/reminders required.	prompts nders used: 0 d them, frequent	r the manuscriporate on this iterate on this iterate of the control of the contro	pt (include quo em by providin our study rs used were prompts ay be necessa	g additional in (letters, emails ry to distinguis	oformation not s, phone calls, sh between the	in the ms, or briefly  SMS) to use the level of
Does your paper Copy and paste relevant sequotes from your manusce explain why the item is not Your answer  6-xi) Report any paper any prompts/reminapplication, what triggered prompts/reminders required.	prompts nders used: 0 d them, frequent	r the manuscriporate on this iterate on this iterate of the control of the contro	pt (include quo em by providin our study rs used were prompts ay be necessa	g additional in (letters, emails ry to distinguis	oformation not s, phone calls, sh between the	in the ms, or briefly  SMS) to use the level of
Does your paper Copy and paste relevant s quotes from your manusc explain why the item is no	prompts nders used: 0 d them, frequent	reminder  Clarify if there verelizability).	pt (include quo em by providin our study  rs used were prompts ay be necessal	g additional in (letters, emails ry to distinguis eminders for a	s, phone calls, sh between the a routine applic	in the ms, or briefly  SMS) to use the level of

# 5-xii) Describe any co-interventions (incl. training/support)

Describe any co-interventions (incl. training/support): Clearly state any interventions that are provided in addition to the targeted eHealth intervention, as ehealth intervention may not be designed as stand-alone intervention. This includes training according and support [1]. It may be proceeded to distinguish between the level of training required for

generalizability.						
	1	2	3	4	5	
subitem not at all important	$\circ$	0	0	0	0	essential

includes training sessions and support [1]. It may be necessary to distinguish between the level of training required for

the trial, and the level of training for a routine application outside of a RCT setting (discuss under item 21 –

#### Does your paper address subitem 5-xii? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Participants were briefly trained to use MyFitnessPal as instructed during the baseline visit. All intervention content was standalone and delivered digitally.

6a) Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed

"The primary outcome was weight change at 3 months... We measured body weight using a calibrated electronic scale (SECA 876) at baseline, 1 month, and 3 months in light clothing with shoes removed. Height was measured to the nearest 0.1 cm using a calibrated, wall-mounted stadiometer (SECA 222). Baseline height was used for calculation of BMI at all time points. We collected self-reported body weight at 6- months and asked participants to send a photo with their feet on the scale displaying the value in either kg or lbs. We assessed the proportion of participants at 3 months who achieved weight loss of  $\geq$ 3% and  $\geq$ 5% from baseline."

"We used a software engine developed at Duke—Prompt—to collect participants' objective MyFitnessPal self-monitoring data; Prompt retrieved this data using the application programming interface (API) of Fitbit, which was linked to participant's MyFitnessPal account. Primary outcomes for self-monitoring engagement span from day 1 (the day after participants' baseline visit) to day 83, and were categorized into the first 4 weeks in the intervention (day 1 to 28), the final two months (day 29-83), and the entire 83-day intervention period. Exploratory analyses examined engagement data after the intervention ended up to 6 months (day 183) post- randomization.

We examined the median number of days per week that participants self-monitored weight, and self-monitored diet (with a complete entry), as well as the percentage of days that entries were recorded (i.e., number of days with entries recorded divided by number of days instructed to record an entry, multiplied by 100). "

# 6a-i) Online questionnaires: describe if they were validated for online use and apply CHERRIES items to describe how the questionnaires were designed/deployed

If outcomes were obtained through online questionnaires, describe if they were validated for online use and apply CHERRIES items to describe how the questionnaires were designed/deployed [9].



#### Does your paper address subitem 6a-i?

Copy and paste relevant sections from manuscript text

Your answer

6a-ii) Describe whether and how "use" (including intensity of use/dosage) was defined/measured/monitored

analysis, etc.). Use/adopt	tion metrics a	are important p	rocess outcor	nes that shou	d be reported i	n any ehealth trial.
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
mportant						
Ooes your paper						
our answer						
\ 5		, ,		1		
5a-iii) Describe v	vhether,	how, and	when qua	alitative fe	edback fr	rom
participants was	obtaine	d				
Describe whether, how, and orms, interviews, focus o		itative feedbac	k from partici	pants was obt	ained (e.g., thro	ough emails, feedba
	1	2	3	4	5	
subitem not at all important	0	0	$\circ$	0	0	essential
Copy and paste relevant solutions	sections fron	n manuscript te	ext			
6b) Any changes	to trial	outcomes	after the	trial com	menced	with
easons	s to trial	outcomes	arter the	tilai con	iiriericeu,	VVICII
Cubbilo						
Does your paper	address	CONSOR	RT subiter	n 6b? *		
Copy and paste relevant						
quotes from your manusc explain why the item is no				ig additional if	normation not	in the ms, or briefly
N/A - no changes we	ere made a	fter trial cor	nmenceme	nt		
7a) How sample	size wa	s determi	ned			
NPT: When applicable,	details of wl	nether and ho	w the clusteri	ing by care pr	ovides or cent	ters was addresse

7a-i) Describe whether and how expected attrition was taken into account

	1	2	3	4	5	
subitem not at all important	$\circ$	0	0	0	0	essential

#### Does your paper address subitem 7a-i?

Copy and paste relevant sections from manuscript title (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Sample size was calculated based on power to detect a 3.5 kg difference in weight change at 3 months between the Sequential arm and the App-Only arm (our primary comparison).[28,29] Our power analysis (G\*Power 3.1.9.2.) determined that 93 participants were needed to achieve 80% power for a two-sided test with an alpha level of 0.05. To account for attrition of 10% and to obtain equal-size groups, we aimed to recruit 105 participants (35 per group)."

# 7b) When applicable, explanation of any interim analyses and stopping guidelines

#### Does your paper address CONSORT subitem 7b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

No interim analyses were conducted. "Participants who became ineligible during the study period up to 3 months were excluded from analyses." Five participants were removed from analyses after they became ineligible.

#### 8a) Method used to generate the random allocation sequence

NPT: When applicable, how care providers were allocated to each trial group

#### Does your paper address CONSORT subitem 8a? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Using simple random sampling, participants were then randomized by study staff to one of three treatment arms using Excel's random number generator to allocate participants equally (1:1:1) across conditions."

8b) Type of randomisation; details of any restriction (such as blocking and block size)

#### Does your paper address CONSORT subitem 8b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Simple random sampling. No blocking/stratification was used.

9) Mechanism used to implement the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assigned

#### Does your paper address CONSORT subitem 9? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Randomization was revealed to participants by study personnel; as such, study staff were not blinded to treatment allocation, but were blinded to the allocation sequence."

10) Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions

#### Does your paper address CONSORT subitem 10? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Using simple random sampling, participants were then randomized by study staff to one of three treatment arms using Excel's random number generator to allocate participants equally (1:1:1) across conditions. Randomization was revealed to participants by study personnel; as such, study staff were not blinded to treatment allocation, but were blinded to the allocation sequence. Participants then reviewed materials describing their treatment condition, in writing and with study staff to reduce contamination."

TIOVV

NPT: Whether or not administering co-interventions were blinded to group assignment

#### 11a-i) Specify who was blinded, and who wasn't

Specify who was blinded, and who wasn't. Usually, in web-based trials it is not possible to blind the participants [1, 3] (this should be clearly acknowledged), but it may be possible to blind outcome assessors, those doing data analysis or those administering co-interventions (if any).

1 2 3 4 5

subitem not at all important O O O O essential

#### Does your paper address subitem 11a-i? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

N/A - it was not possible to blind study staff nor participants to treatment condition.

# 11a-ii) Discuss e.g., whether participants knew which intervention was the "intervention of interest" and which one was the "comparator"

Informed consent procedures (4a-ii) can create biases and certain expectations - discuss e.g., whether participants knew which intervention was the "intervention of interest" and which one was the "comparator".

1 2 3 4 5

subitem not at all important O O O essential

## Does your paper address subitem 11a-ii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

# 11b) If relevant, description of the similarity of interventions

(this item is usually not relevant for ehealth trials as it refers to similarity of a placebo or sham intervention to a active medication/intervention)

#### Does your paper address CONSORT subitem 11b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the providing why the item is not applicable (relevant for your study).

All three arms shared common components, described below. The "App-Only" arm added no additional content. The two more enhanced intervention arms added feedback and other intervention content, and also differed in the timing of delivery of different intervention components (Simultaneous vs Sequential tracking of behaviors).

"All treatment arms self-monitored dietary intake using MyFitnessPal, a free commercial app that allows users to log food and beverages, and provides nutritional information from a database with over 5 million foods (my tnesspal.com). In-app feedback in both graphical and text format provides users with real-time progress updates. Participants received a goal to lose 5% of their initial body weight by 12 weeks. Based on this goal and the participant's current weight, a weekly weight loss goal between 0.5 and 2.0 pounds was calculated. Along with the Mifflin-St. Jeor equation that factors in basal metabolic rate, this value was used to determine a tailored daily calorie goal, with a minimum caloric goal of 1200 kcal/day for women and 1500 kcal/day for men. During the baseline visit, in-app push-reminders were programmed to be sent each day if tracking had not occurred by a pre-specified time. No structured dietary advice (e.g., follow a low carbohydrate diet) was given to participants. In the App-Only arm, MyFitnessPal served as an "off- the-shelf," self-guided approach that the general U.S. population can already access for free in the commercial marketplace."

# 12a) Statistical methods used to compare groups for primary and secondary outcomes

NPT: When applicable, details of whether and how the clustering by care providers or centers was addressed

"We used intent-to-treat (ITT) analyses to test our primary aim using linear mixed modeling with an unstructured covariance matrix and restricted maximum likelihood estimates to examine changes in weight over time by treatment arm. We assumed missing at random and used SAS 9.4 PROC MIXED (SAS Institute, Cary, NC) for these analyses. For 6-month weight values sent via photo, we subtracted 0.172 kg (.4 lbs) to account for participants holding a device on the scale to take the photo. To account for the 6-month self-reported weight data without photos, we used a regression model to adjust for age, gender, and race/ethnicity.[30] Participants who sent a photo of their 6-month weight did not differ on any measured sociodemographic characteristics from those who did not send a photo (data not shown). We used chi-square tests to assess proportion of participants achieving ≥3% and ≥5% weight loss; we assumed non-completers did not achieve this clinical threshold."

#### 12a-i) Imputation techniques to deal with attrition / missing values

Imputation techniques to deal with attrition / missing values: Not all participants will use the intervention/comparator as intended and attrition is typically high in ehealth trials. Specify how participants who did not use the application or dropped out from the trial were treated in the statistical analysis (a complete case analysis is strongly discouraged, and simple imputation techniques such as LOCF may also be problematic [4]).

	1	2	3	4	5	
subitem not at all important	$\circ$	$\circ$	0	0	0	essential

#### Does your paper address subitem 12a-i? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"We assumed missing at random "

"We used chi-square tests to assess proportion of participants achieving ≥3% and ≥5% weight loss; we assumed non-completers did not achieve this clinical threshold."

12b) Methods for additional analyses, such as subgroup analyses and adjusted analyses

## Does your paper address CONSORT subitem 12b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the providing additional information of the ms, or briefly available to the providing additional information of the ms, or briefly available to the providing additional information of the ms, or briefly available to the manuscript of the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the manuscript of the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the manuscript of the

"For the baseline characteristics, we computed descriptive statistics, stratified by treatment arm or retention status. We used the Pearson chi-square test for categorical variables, analysis of variance for continuous variables, and Fisher's exact tests with small cell counts. All analyses were two-tailed. Participants who became ineligible during the study period up to 3 months were excluded from analyses."

"Given non-normally distributed intervention engagement data, we reported medians and interquartile ranges (IQR). To examine differences between treatment arms, we used Wilcoxon Mann-Whitney U tests (if two arms) and the Kruskal-Wallis tests (if three arms). We used Spearman rank correlation coefficients (rs) to examine the relation between self-monitoring engagement and change in weight. We also assessed for contamination by exploring whether participants self-monitored when they were not expected to do so."

# X26) REB/IRB Approval and Ethical Considerations [recommended as subheading under "Methods"] (not a CONSORT item)

#### X26-i) Comment on ethics committee approval

	1	2	3	4	5	
subitem not at all important	$\circ$	$\circ$	0	$\circ$	0	essential

#### Does your paper address subitem X26-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

# x26-ii) Outline informed consent procedures

Outline informed consent procedures e.g., if consent was obtained offline or online (how? Checkbox, etc.?), and what information was provided (see 4a-ii). See [6] for some items to be included in informed consent documents.

	1	2	3	4	5	
subitem not at all important	$\circ$	0	0	0	0	essential

#### Does your paper address subitem X26-ii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms and the providing additional information not in the ms.

#### X26-iii) Safety and security procedures

Safety and security procedures, incl. privacy considerations, and any steps taken to reduce the likelihood or detection of harm (e.g., education and training, availability of a hotline)

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

#### Does your paper address subitem X26-iii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

#### **RESULTS**

13a) For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome

NPT: The number of care providers or centers performing the intervention in each group and the number of patients treated by each care provider in each center

## Does your paper address CONSORT subitem 13a? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

See CONSORT diagram (Figure 1)

13b) For each group, losses and exclusions after randomisation, together with reasons

Does your paper address CONSORT subitem 13b? (NOTE: Preferably, this is shown in a CONSORT flow diagram) \*

quotes from your manusc explain why the item is no				g additional ir	nformation not ir	n the ms, or briefly
See CONSORT diagra						
13b-i) Attrition d	iagram					
Strongly recommended: A intervention/comparator demonstrating usage/dos	in each group	plotted over t				
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper					formula de la Cara	hada washa in
Copy and paste relevant s quotation marks "like this additional information no	" to indicate	direct quotes f	rom your man	iscript), or ela	borate on this it	em by providing
see Figure 4 (self-mo	onitoring p	er interventi	on week by	treatment a	arm)	
14a) Dates defin	ing the p	periods of	recruitme	ent and fo	ollow-up	
Does your paper Copy and paste relevant s					on marke "like th	aie" to indicate direct
quotes from your manusc explain why the item is no	cript), or elab	orate on this it	em by providin	•		
"Recruitment occurre			-	ı		
14a-i) Indicate if	critical '	ʻsecular e	vents" fel	into the	study perio	od
Indicate if critical "secular "changes in computer har	r events" fell	into the study	period, e.g., sig			
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

## Does your paper address subitem 14a-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the providing additional information and in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information in the ms, or briefly available to the providing additional information and the providing a

#### 14b) Why the trial ended or was stopped (early)

#### Does your paper address CONSORT subitem 14b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

N/A - trial was completed as intended

# 15) A table showing baseline demographic and clinical characteristics for each group

NPT: When applicable, a description of care providers (case volume, qualification, expertise, etc.) and centers (volume) in each group

#### Does your paper address CONSORT subitem 15? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Yes - table 4

## 15-i) Report demographics associated with digital divide issues

In ehealth trials it is particularly important to report demographics associated with digital divide issues, such as age, education, gender, social-economic status, computer/Internet/ehealth literacy of the participants, if known.

	1	2	3	4	5	
subitem not at all important	$\circ$	0	$\circ$	0	0	essential

## Does your paper address subitem 15-i? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Yes - table 4

# 16-i) Report multiple "denominators" and provide definitions Report multiple "denominators" and provide definitions: Report N's (and effect sizes) "across a range of study participation [and use] thresholds" [1], e.g., N exposed, N consented, N used more than x times, N used more than y weeks, N participants "used" the intervention/comparator at specific pre-defined time points of interest (in absolute and relative numbers per group). Always clearly define "use" of the intervention. 1 2 3 4 5 subitem not at all important O O O O essential

#### Does your paper address subitem 16-i? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Yes - see tables

#### 16-ii) Primary analysis should be intent-to-treat

Primary analysis should be intent-to-treat, secondary analyses could include comparing only "users", with the appropriate caveats that this is no longer a randomized sample (see 18-i).

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

#### Does your paper address subitem 16-ii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"We used intent-to-treat (ITT) analyses to test our primary aim using linear mixed modeling with an unstructured covariance matrix and restricted maximum likelihood estimates to examine changes in weight over time by treatment arm."

17a) For each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)

## Does your paper address CONSORT subitem 17a? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms and the providing additional information not in the ms.

# 17a-i) Presentation of process outcomes such as metrics of use and intensity of use

In addition to primary/secondary (clinical) outcomes, the presentation of process outcomes such as metrics of use and intensity of use (dose, exposure) and their operational definitions is critical. This does not only refer to metrics of attrition (13-b) (often a binary variable), but also to more continuous exposure metrics such as "average session length". These must be accompanied by a technical description how a metric like a "session" is defined (e.g., timeout after idle time) [1] (report under item 6a).

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

#### Does your paper address subitem 17a-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Table 6

# 17b) For binary outcomes, presentation of both absolute and relative effect sizes is recommended

#### Does your paper address CONSORT subitem 17b? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Yes, in text and more detail in tables. For example:

"The proportion of participants achieving at least 3% weight loss at 3 months was similar between arms (P=.43; Simultaneous: 41% [13 of 32]; Sequential: 44% [13 of 34]; App-Only: 29% [10 of 34]). Likewise, weight loss of at least 5% at 3-months did not significantly differ among arms (P=.26) and occurred in 31% (10 of 32), 21% (7 of 34), and 15% (5 of 34), respectively."

18) Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory

#### Does your paper address CONSORT subitem 18?\*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the item is not applicable trajectory to trudy.

"Contamination. The median [IQR] frequency of days that App-Only participants tracked weight during the 3-month intervention was 1% [8%], and the frequency of days that Sequential participants tracked diet during month 1 was 0% [0%] (see Table 6 for absolute values)."

#### 18-i) Subgroup analysis of comparing only users

A subgroup analysis of comparing only users is not uncommon in ehealth trials, but if done, it must be stressed that this is a self-selected sample and no longer an unbiased sample from a randomized trial (see 16-iii).

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

#### Does your paper address subitem 18-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

#### 19) All important harms or unintended effects in each group

(for specific guidance see CONSORT for harms)

#### Does your paper address CONSORT subitem 19? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

N/A - there were no unintended effects

## 19-i) Include privacy breaches, technical problems

Include privacy breaches, technical problems. This does not only include physical "harm" to participants, but also incidents such as perceived or real privacy breaches [1], technical problems, and other unexpected/unintended incidents. "Unintended effects" also includes unintended positive effects [2].

	1	2	3	4	5	
subitem not at all important	$\circ$	$\circ$	0	0	0	essential

#### Does your paper address subitem 19-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms, or briefly available to the providing additional information not in the ms.

# 19-ii) Include qualitative feedback from participants or observations from staff/researchers

Include qualitative feedback from participants or observations from staff/researchers, if available, on strengths and shortcomings of the application, especially if they point to unintended/unexpected effects or uses. This includes (if available) reasons for why people did or did not use the application as intended by the developers.

	1	2	3	4	5	
subitem not at all important	$\circ$	0	0	0	0	essential

#### Does your paper address subitem 19-ii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

#### **DISCUSSION**

22) Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence

NPT: In addition, take into account the choice of the comparator, lack of or partial blinding, and unequal expertise of care providers or centers in each group

# 22-i) Restate study questions and summarize the answers suggested by the data, starting with primary outcomes and process outcomes (use)

Restate study questions and summarize the answers suggested by the data, starting with primary outcomes and process outcomes (use).

	1	2	3	4	5	
subitem not at all important	$\circ$	$\circ$	0	0	0	essential

#### Does your paper address subitem 22-i? \*

"A low-intensity intervention utilizing a commercial app for self-monitoring resulted in comparable weight loss at 3 months, with no variability by treatment arm. Loss of 3-5% of initial weight has been linked to improved health outcomes,[31-32] suggesting that GoalTracker is an efficacious intervention for clinically-meaningful weight loss. GoalTracker's Simultaneous arm had a comparable or higher proportion of participants achieving 5% weight loss, compared to other weight loss interventions that used mobile apps for self-monitoring dietary intake (range of 26- 35%;[16,33-35]), but lower rates than some interventions including counseling (range of 42-44%;[17,36-37]). Interestingly, we found sustained weight loss at 6 months, with trends suggesting continued weight loss in the Simultaneous arm, relative to weight gain in the other two arms."

#### 22-ii) Highlight unanswered new questions, suggest future research

Highlight unanswered new questions, suggest future research.

1 2 3 4 5

subitem not at all important O O O o essential

#### Does your paper address subitem 22-ii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

# 20) Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses

## 20-i) Typical limitations in ehealth trials

Typical limitations in ehealth trials: Participants in ehealth trials are rarely blinded. Ehealth trials often look at a multiplicity of outcomes, increasing risk for a Type I error. Discuss biases due to non-use of the intervention/usability issues, biases through informed consent procedures, unexpected events.

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

#### Does your paper address subitem 20-i? \*

"Because this study was powered on superiority rather than equivalency, we cannot definitively assert that the treatment arms produce comparable weight loss. In addition, we collected self-reported weight at 6-months due to logistical reasons; however, we were encouraged to find that no additional attrition occurred between 3 and 6 months, despite no contact occurring during that period and no incentive given to provide a weight value. Additionally, neither study staff nor participants were blinded to treatment arm, and we required participants to have access to a bathroom scale, though this mimics the real-world population who would track weight. Lastly, this study did not include a pure control arm without an intervention, which may have led to an underestimation of treatment effects, as could the possibility of data not actually missing at random."

#### 21) Generalisability (external validity, applicability) of the trial findings

NPT: External validity of the trial findings according to the intervention, comparators, patients, and care providers or centers involved in the trial

#### 21-i) Generalizability to other populations

Generalizability to other populations: In particular, discuss generalizability to a general Internet population, outside of a RCT setting, and general patient population, including applicability of the study results for other organizations

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

#### Does your paper address subitem 21-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

# 21-ii) Discuss if there were elements in the RCT that would be different in a routine application setting

prompts/reminders, more omission of these elements setting.						
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address subitem 21-ii?  Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study  Your answer						
OTHER INFORM	ATION	7				
					,	
23) Registration	number	and name	e of trial r	egistry		
Does your paper	address	CONSOF	RT subiter	n 23? *		

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Clinical Trial Registration: clinicaltrials.gov; identifier NCT03254953"

## 24) Where the full trial protocol can be accessed, if available

#### Does your paper address CONSORT subitem 24? \*

Cite a Multimedia Appendix, other reference, or copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

https://clinicaltrials.gov/ct2/show/NCT03254953

# 25) Sources of funding and other support (such as supply of drugs), role of funders

#### Does your paper address CONSORT subitem 25? \*

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly available to the item is not applicable trajected for your study.

"The research described in this paper was supported by a grant to the first author from the American Psychological Association, the Duke Interdisciplinary Behavioral Research Center, and the Aleane Webb Dissertation Research Award provided by The Graduate School at Duke University."

#### X27) Conflicts of Interest (not a CONSORT item)

# X27-i) State the relation of the study team towards the system being evaluated

In addition to the usual declaration of interests (financial or otherwise), also state the relation of the study team towards the system being evaluated, i.e., state if the authors/evaluators are distinct from or identical with the developers/sponsors of the intervention.

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

#### Does your paper address subitem X27-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

#### About the CONSORT EHEALTH checklist

As a result of using this checklist, did you make changes in your manuscript?

^	
0	yes, major changes
•	yes, minor changes
0	no

What were the most important changes you made as a result of using this checklist?

How much time did you spend on going through the checklist INCLUDING making changes in your manuscript *
3.5 hours
As a result of using this checklist, do you think your manuscript has improved? *
yes
O no
Other:
Would you like to become involved in the CONSORT EHEALTH group?  This would involve for example becoming involved in participating in a workshop and writing an "Explanation and Elaboration" document
O yes
O no
Other:
Any other comments or questions on CONSORT EHEALTH
Your answer

# STOP - Save this form as PDF before you click submit

To generate a record that you filled in this form, we recommend to generate a PDF of this page (on a Mac, simply select "print" and then select "print as PDF") before you submit it.

When you submit your (revised) paper to JMIR, please upload the PDF as supplementary file.

Don't worry if some text in the textboxes is cut off, as we still have the complete information in our database. Thank you!

# Final step: Click submit!

Click submit so we have your answers in our database!



Never submit passwords through Google Forms.

 $This \ content \ is \ neither \ created \ nor \ endorsed \ by \ Google. \ Report \ Abuse - Terms \ of \ Service - \ Additional \ Terms$ 

Google Forms